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Long-chain GM1 gangliosides alter transmembrane domain registration through interdigitation

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Abstract

Extracellular and cytosolic leaflets in cellular membranes are distinctly different in lipid composition, yet they contribute together to signaling across the membranes. Here we consider a mechanism based on long-chain gangliosides for coupling the extracellular and cytosolic membrane leaflets together. Based on atomistic molecular dynamics simulations, we find that long-chain GM1 in the extracellular leaflet exhibits a strong tendency to protrude into the opposing bilayer leaflet. This interdigitation modulates the order in the cytosolic monolayer and thereby strengthens the interaction and coupling across a membrane. Coarse-grained simulations probing longer time scales in large membrane systems indicate that GM1 in the extracellular leaflet modulates the phase behavior in the cytosolic monolayer. While short-chain GM1 maintains phase-symmetric bilayers with a strong membrane registration effect, the situation is altered with long-chain GM1. Here, the significant interdigitation induced by long-chain GM1 modulates the behavior in the cytosolic GM1-free leaflet, weakening and slowing down the membrane registration process. The observed physical interaction mechanism provides a possible means to mediate or foster transmembrane communication associated with signal transduction.

Key words:

Glycosphingolipid, cholesterol, membrane domain, membrane registry, molecular dynamics, computer simulations

Introduction

Cells invest substantial resources to synthesize a broad range of membrane lipids and to maintain their asymmetric inhomogeneous distributions in biological membranes [1, 2]. In addition to roles in maintaining membrane stability and fluidity, awareness is growing that the headgroups of glycosphingolipids (GSLs) are of particular relevance for recognition processes, especially by tissue lectins [3-5]. Notably the presence of glycolipids and the mode of presentation are of critical significance to serve as functional counterreceptors [6]. In this respect, organization of lipids as in-plane membrane domains has been studied for decades [7-9], yet an understanding of domain structure and composition, and the biological function of domains overall, is still quite incomplete. The concept of lipid rafts [10-12] (functional nanoscale domains rich in cholesterol, saturated phospholipids, and bioactive lipids such as gangliosides) as well as different interpretations of research data [13] concur with the existence of functional membrane units comprised of proteins and lipids, where lipid-protein interactions modulate several aspects of protein activity [14]. This suggests that membrane-associated proteins can reorganize their membrane environment to be rich in lipids that modulate protein features dynamically. In comparison, under protein-free conditions the existence of membrane domains with different lipid compositions and physical properties is well established [15]. Studies on model membranes have unambiguously shown macroscopic phase separation into cholesterol-rich liquid-ordered (L_o) and cholesterol-poor liquid-disordered (L_d) domains in a manner [16, 17], where the phase separation in the bilayer plane arises from lipid-dependent liquid-liquid immiscibility.

Due to their already noted asymmetry, the two leaflets of biological membranes are also different in composition [1, 2]. The lipid mixture that is typical for the extracellular leaflet of eukaryotic plasma membranes contains sphingolipids with a high percentage of cholesterol. It tends to phase separate when reconstituted in model membranes. In contrast, such phase separation does not take place for the lipid mixture that represents the cytoplasmic leaflet enriched in, e.g., unsaturated lipids [18, 19].

The distinctly different transmembrane compositions of biological membranes can lead to phase asymmetry or antiregistration, meaning that L_o and L_d domains overlap each other in the two opposite leaflets [20,21]. Meanwhile, a domain in one leaflet can induce formation of a similar domain also in the opposite leaflet (i.e., both being in the L_o or L_d phase), leading to phase symmetry or registration [20-22]. Several mechanisms for protein-independent domain induction between membrane leaflets have been proposed, including

interfacial energy minimization [23], electrostatic coupling [20], lipid and cholesterol flip-flop [20, 24], composition-curvature coupling [25, 26], and dynamic chain interdigitation [20, 27, 28]. However, despite considerable research efforts, there is no consensus on whether biological membranes are phase symmetric or asymmetric. Furthermore, the exact mechanism of induced domain formation remains undetermined.

There is evidence in favor of both bilayer registration and antiregistration [18, 20, 23, 29] but the overall understanding of their causes is still unclear. This partly stems from difficulties to carry out robust and unambiguously interpretable experiments on domains in biological membranes, where, for instance, one applies methods such as detergent extraction that is unable to characterize the phases in individual bilayer leaflets. On the other hand, experiments on asymmetric model membranes are carried out under non-equilibrium conditions, right after the formation of an asymmetric membrane, since lipid flip-flops quite rapidly lead to transbilayer symmetry. Although these studies have significantly enhanced our current understanding of this topic, the interpretation of experimental data is admittedly not straightforward.

Theoretical and simulation studies provide another means to explore the coupling between membrane domains in different leaflets and to elucidate the physical principles that lead to domain registration and/or antiregistration [20, 26, 30-33]. These studies have suggested that the coupling of phase behavior between the leaflets is highly sensitive to lipid composition, yet it is not well understood why certain lipid mixtures induce domain registration whereas others do not.

Based on the concept of the sugar code that ascribes informational contents to glycans presented by scaffolds (sphingolipids, proteins) [34-36], giving special attention to GSLs is warranted. GSLs are integral components of ordered lipid domains [37, 38], and their carbohydrate headgroup is linked to a ceramide moiety, exclusively sorted to the extracellular leaflet of plasma membranes, where they can engage in biorecognition [39, 40]. We here focus on ganglioside GM1, a so-called true factotum of nature [41], because it is a counterreceptor for adhesion/growth-regulatory galectins, hereby involved for example in effector/regulatory T cell communication, lectin endocytosis, neuritogenesis, and neuroblastoma growth arrest [42-47]. To efficiently do so, positional aspects (density and presentation) appear crucial [48]. Notably, the interaction with the human bioeffector has also been analyzed in model membranes without proteins [49]. Looking at the structure of GSLs, a source of variability concerns the length of the acyl chain. While the majority of phospholipids have acyl chains with 16 to 18 carbons, GSLs can typically have considerably

longer chains that consist of 24 carbons or more [50-52], allowing GSLs to pack tightly into ordered membrane domains. The long hydrocarbon chains may promote greater exposure of the carbohydrate headgroup and therefore enhance ligand binding of GSLs [51]. In fact, the routing signal for apical and axonal transport of glycoproteins by galectin-4, i.e. the sulfatide headgroup, is presented preferentially by ceramide with long (C24)-chain fatty acids [52-54]. As a further consequence, a long tail can extend into the opposite bilayer leaflet through interdigitation and thereby provide a potential means for signal transduction [50]. Given that contribution by both modes of placement of the long acyl chain can be of physiological relevance, consideration of the effect of GSL chain length on membrane registration is of interest.

To address this issue, we carried out a series of atomistic and coarse-grained molecular dynamics (MD) simulations on many-component membranes with GM1 to unravel how its acyl chain length (ranging from 16 to 30 carbons) contributes to membrane registration. We found that long-chain GM1 interdigitates to a substantial degree to the opposite membrane leaflet and alters its phase behavior.

Methods

Atomistic simulations of symmetric bilayers

We used all-atom MD simulations to consider eight types of model membranes that were composed of 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC) (202 molecules), GM1 (14 molecules), and cholesterol (Chol) (72 molecules). These eight systems were also simulated in the absence of cholesterol. The molar concentrations of GM1 (POPC) were therefore 4.9 (70.1) and 6.5 (93.5) mol% in systems with and without cholesterol, respectively. Cholesterol concentration was 25 mol% in the three-component system (Table 1). The transmembrane lipid distributions were in all systems symmetric. The factor that differentiates the eight types of bilayers is the length of the fatty acid acyl chain of GM1: it ranged from palmitic acid (16:0, referred to as GM1₁₆) to melissic acid (30:0, referred to as GM1₃₀). Given the abundance of both POPC and cholesterol, the POPC/cholesterol/GM1 mixture provides an appropriate basis to consider the effects of GM1 in a system, whose phase behavior has been suggested to be rich, especially when it is extended with other lipid components (see, e.g., [55]). Figure 1 shows the chemical structure of all lipid molecules

involved in this study. Bilayers were extensively hydrated with 13785 water molecules, and 14 Na⁺ counterions were added to neutralize the excess charge of GM1.

Lipid molecules were described by the all-atom OPLS force field [56] with recent extensions for carbohydrates [57] and lipids [58-61]. Water was described by the TIP3P model [62] that is compatible with the OPLS-AA force field. Prior to MD simulations the systems were energy-minimized using the steepest-descend algorithm. For each of the 16 systems, an MD simulation of 400 ns was conducted using the GROMACS 4.6.6 software package [63]. The first 100 ns period of the trajectories was considered as equilibration and the last 300 ns of the trajectories were used for analyses.

Simulations were performed in the isobaric-isothermal (NpT) ensemble (310 K and 1 bar). The temperature of the system was maintained by the velocity-rescaling (v-rescale) thermostat with a time constant of 0.1 ps [64]. The temperatures of the solute and the solvent were controlled independently. The pressure of the system was maintained semiisotropically by the Parrinello–Rahman barostat with a 1 ps time constant [65]. The time step for integration of equations of motion was set to 2 fs. Periodic boundary conditions were imposed in all three dimensions. For the long-range electrostatic interactions, the particle-mesh Ewald (PME) method was used [66]. Lennard–Jones interactions were cutoff at a distance of 1.0 nm. The linear constraint solver (LINCS) algorithm was used to preserve covalent bond lengths [67].

Coarse-grained simulations of asymmetric bilayers

To study the effect of acyl chain length of GM1 on membrane phase behavior, we employed the coarse-grained Martini model [68-70]. A bilayer consisting of 1348 lipids (674 in each leaflet) was first constructed as a random arrangement of 30 mol% palmitoylsphingomyelin (PSM), 40 mol% dilinoleylphosphatidylcholine (DLiPC), and 30 mol% cholesterol (Figure 1). This system was then used for preparing four systems with varying concentration and acyl chain length of GM1 (Table 1). First, either 1.5 or 6.0 mol% of GM1 was inserted into the upper leaflet by randomly replacing existing lipids from this leaflet with GM1 in a manner where the 30:40:30 ratio between PSM, DLiPC, and cholesterol, respectively, was maintained. Second, the fatty acid attached to these GM1 molecules was chosen to be either five (referred to as GM1_{short}) or eight beads (referred to as GM1_{extended}) long (Figure 1). The “short” and “extended” cases correspond to GM1 molecules with an acyl chain of ~18 and ~30 carbons, respectively. The ternary DLiPC/PSM/cholesterol system chosen for coarse-grained simulations is largely the standard system employed in phase separation studies

using the Martini approach (see, e.g., refs. [71,72]). The systems were hydrated with 18110 water beads. About 150 mM of NaCl was added to the system together with counterions to compensate for the charge of GM1s.

The four GM1-containing bilayers were simulated for 10 μ s each, using a time step of 5 fs. We performed 10 repeats for each bilayer (4 x 10 x 10 μ s) (Table 1). A GM1-free control system (as a symmetric bilayer) was simulated for 10 μ s with a time step of 25 fs (Table 1). For the GM1-free control system, we performed 8 repeats. The temperature of the lipids and the solvent were separately maintained at 300 K using the velocity-rescaling thermostat with a time constant of 1 ps [64]. Pressure was coupled semiisotropically to the Parrinello–Rahman barostat [65]. A reference pressure of 1 bar was employed, and the coupling time was set to 12 ps. The Verlet cutoff scheme was employed for nonbonded interactions. Electrostatic interactions were handled with the reaction field method with a cut-off of 1.1 nm and the relative permittivity of 15. The potential shift modifier was employed for the Lennard–Jones interactions and a cut-off of 1.1 nm was used. Periodic boundary conditions were employed in all three dimensions. All coarse-grained simulations were run with GROMACS 5.0.x [73].

Analyses

Area per lipid was calculated by dividing the time-averaged total area of the simulation box (in the bilayer plane) by the number of lipids in a single leaflet. Bilayer thickness was computed from the distance between average positions of phosphate atoms in the two bilayer leaflets. The deuterium order parameter S_{CD} describing the conformational order along lipid acyl chains is defined for each carbon in the chain as in [74]:

$$S_{CD,i} = \left\langle \frac{3}{2} \cos^2 \theta_i - \frac{1}{2} \right\rangle,$$

where θ_i is the angle between a C–D bond (C–H in simulations) for carbon number i and the bilayer normal. The angular brackets denote averaging over time and lipids of the same type.

To characterize the conformation of GM1’s pentasaccharide, we analyzed the orientation of the GM1 headgroup in terms of three vectors depicted in Figure S1. The first vector was named as *Wrist* based on its location in the GM1 headgroup, describing the orientation of the first sugar (Glc) that is O-glycosidically connected to ceramide. The two

other vectors (*Thumb* and *Forefinger*) characterize the orientations of the two outermost branches in the GM1 head group (Figure S1) that are the contact sites for human and bacterial lectins [75]. The angles between these vectors and the bilayer normal provide an overall picture of the head group conformation.

Interdigitation between the two membrane leaflets is characterized by the overlap parameter $q_{oi}(z)$ as [76]

$$q_{oi}(z) = 4 \frac{q_i(z)q_o(z)}{[q_i(z)+q_o(z)]^2},$$

where z is the coordinate in the membrane normal direction, and $q_i(z)$ and $q_o(z)$ are the number densities of atoms belonging to the inner and outer leaflets, respectively, calculated from a trajectory centered at the middle of a bilayer center. When the overlap parameter is integrated over the membrane along the bilayer normal direction (over the simulation box from zero to d), one finds the length scale

$$\lambda = \int_0^d q_{oi}(z) dz ,$$

where λ describes in a suggestive fashion the length scale over which the two monolayers are interdigitated. Here, one has to keep in mind that λ does not describe the extent of interdigitation of individual lipid chains. Rather, it provides a measure for the average interdigitation between the two leaflets.

In CG simulations of the many-component membranes, we quantified the extent of in-plane phase separation by the contact fraction between saturated (PSM) and unsaturated (DLiPC) lipids using the approach of Domanski et al. [77],

$$f_{\text{mix}} = \frac{c_{\text{DLiPC-PSM}}}{c_{\text{DLiPC-PSM}} + c_{\text{DLiPC-DLiPC}}},$$

where $c_{\text{DLiPC-PSM}}$ denotes contacts between the phosphate (PO4) beads of DLiPC and PSM molecules. Here, a cut-off distance of 1.1 nm was employed for the determination of contacts. The contact fraction f_{mix} was computed only within the GM1-free leaflet of the asymmetric CG membrane. In a randomly distributed leaflet with no phase separation, f_{mix} is

large (e.g., about 0.41 in the CG system with 6 mol% GM1), but decreases the more the system phase separates.

The rate of phase separation was quantified by fitting the time evolution of the contact fractions (f_{mix}) with an exponentially decaying function

$$f_{\text{mix}}(t) = Ae^{-t/\tau} + b ,$$

where A and b are constants related to the initial and final levels of separation, t is the simulation time, and τ is a decay constant associated with the phase separation process.

In CG model systems, where phase separation into L_o and L_d domains was observed, we analyzed the transbilayer domain registration of regions with a similar degree of order, *i.e.* the membrane registry effect. To this end, for every analyzed frame, the saturated (PSM) and unsaturated (DLiPC) lipids were binned onto a 2D grid based on the position of the phosphate bead (PO4). The cell area was set to 1 nm² and the binning was performed every 10 ns. Then, we checked whether the corresponding cells in the 2D grids of the two leaflets were mostly occupied by the same lipid type (saturated vs. unsaturated). If this condition was satisfied, transbilayer domain registration for this cell was registered. The number of cells with observed coupling was then counted for each studied frame, and the data were normalized so that the value of one stands for perfect registry, where an L_o domain of one leaflet faces an L_o domain of the opposing leaflet, and similarly an L_d domain faces an L_d domain of the opposing leaflet. Similarly, the value of zero stands for perfect antiregistration, where an L_o domain in one leaflet faces an L_d domain of the opposing leaflet and vice-versa. Intermediate values describe the fraction of the system that is in registry.

In consideration of lateral diffusion, where the aim was to determine whether GM1 interdigitation affects the fluidity in the opposing leaflet (see the results below), the mean-squared displacement (MSD) of lipids was calculated from the first 2 μ s, *i.e.*, from the region in time where phase separation takes place. The fits to linear behavior in time were performed to the MSD data between 20–200 ns lag times.

Results

Long acyl chains of GM1 induce interleaflet coupling and ordering that span the whole membrane

Snapshots of atomistic simulations (Figure 2) clearly show the tendency of long acyl chains of GM1 to extend deep into the opposing bilayer leaflet. There is a significant reorganization of hydrocarbon chains at the midplane of the bilayer as demonstrated by the density profiles of the long acyl chains depicting a peak in the bilayer center, in contrast to a dip observed with short-tail GM1 (see Figure S2 in Supplementary Information).

To quantify interdigitation in more detail, Figure S3 shows the partial densities of the terminal carbon atoms of the acyl and sphingosine tails of GM1₁₆ and GM1₃₀, as well as of the terminal carbon of the POPC *sn*-2 chain. The results for GM1₃₀ are quite revealing. The terminal carbon of the GM1 acyl chain protrudes about 2 nm into the opposite layer. This means that GM1₃₀ penetrates *through* the membrane to the head group region in the opposing membrane leaflet. The distribution of GM1₃₀ also has two maxima: the first maximum is located in the middle of the bilayer, characterizing acyl chains that lie along the membrane plane in the free volume cavity, and the second maximum resides deep in the opposite leaflet about 1.5 nm from the bilayer center. Cholesterol strengthens interdigitation, but the interdigitation is exceptionally pronounced without cholesterol, too. The two quite different orientations are also evident from the simulation snapshots in Figure 2. Meanwhile, the terminal carbons of GM1₁₆, the GM1 sphingosine chains, and the POPC *sn*-2 chain behave regularly.

The threshold length of GM1 to observe the double-peak structure in the terminal carbon distribution is about 24 (Figure 3). For shorter acyl chains, calculations for GM1 result in a single peak around the membrane center. For chain lengths of 24 and longer, the double-peak structure arises and becomes more and more prominent for increasing acyl chain length. Cholesterol increases the effect.

Figure 4 highlights the above-discussed findings in terms of λ , which characterizes the length scale over which the two leaflets come into contact to a significant degree. The interdigitation becomes stronger for increasing GM1 acyl chain length. As to the role of cholesterol, GM1₃₀ in the absence of cholesterol results in the same value for λ as GM1₁₆ in the presence of cholesterol.

Interdigitation of long acyl chains of GM1 has a quite dramatic effect on the acyl chain conformational order described by S_{CD} (Figure 5). When the length of the acyl chains rises above ~24, the order parameter profile first levels off and then starts to increase again, reflecting how the terminal region of the GM1 acyl chain aligns with the hydrocarbon chains in the opposing leaflet. The effect is exceptionally strong with cholesterol. Similar behavior

with long acyl chains was recently observed for sphingomyelin in atomistic MD simulations [27], and with long free fatty acids in both MD simulations and NMR measurements [78]. In older studies, a similar shape of the S_{CD} profile was observed for highly asymmetric phosphatidylcholines [79]. In this work, we found that apart from the GM1 acyl chain, the interdigitation does not cause notable changes to the properties of other lipid chains, such as the sphingosine chain of GM1 or the acyl chains of POPC.

Atomistic simulations also revealed that the physical lipid bilayer properties such as average area per lipid and membrane thickness were not affected to any significant degree by changes in acyl chain length of GM1. Further, we found that the length of the acyl chain did not influence the conformation behavior and the orientation of the GM1 head group. These results (Figure S1, Figure S4, Table S2) are presented in Supplementary Information.

Long-chain GM1 perturbs the phase behavior of the opposing leaflet

The above atomistic results suggest that the interdigitation of long-chain GM1 such as GM1₃₀ reaches a level that can alter the order and thereby also the physical phase of the opposing leaflet. Large-scale simulations of the CG models were performed to test this assumption. Figure 6 shows structures for asymmetric many-component GM1:PSM:DLiPC:Chol membranes, where GM1 gangliosides with an extended acyl tail (GM1_{extended}) in the upper leaflet interdigitate to the lower monolayer. The membrane system in question undergoes spontaneous phase separation into L_o and L_d phases in the absence as well as in the presence of GM1. GM1 partitions to the L_o -phase, as expected, and it does not alter the in-plane phase behavior of the leaflet that hosts GM1 (Figure 6). However, the simulation structures (Figure 6) suggest that the long GM1_{extended} chains perturb the structure of the opposite leaflet. More importantly, the data also propose that the transbilayer domain registry is altered by GM1_{extended}. To explore this idea, we first monitored the phase separation of the GM1-free leaflet in time. Figure 7 depicts for the GM1-free membrane that the contact fraction f_{mix} decreases to a significant degree in the course of time, indicating considerable phase separation. The systems with GM1_{short} result in largely the same view. However, with long-chain GM1_{extended} the situation changes. The long-chain GM1 perturbs the opposite leaflet to an extent that also influences its phase behavior: the phase separation process is slowed down, and the extent of phase separation is weakened compared to systems with GM1_{short} or without any GM1. However, the diffusion coefficients of lipids in the opposite leaflet remain unaffected by the GM1 acyl chain length (Figure S5) and consequently the observed slowing down of phase separation does not result from reduced

mobility of lipids.

The rate of phase separation was quantified by fitting the time course of the contact fractions with an exponentially decaying function (see Methods). The average value of the decay constants (determined from the ten replicas for each system) was found to be 0.7 and 0.8 μs with 1.5 and 6 mol% of GM1_{short}, respectively. With long-chain GM1, the average decay constant was observed to be 0.7 μs with 1.5 mol% and 0.9 μs with 6 mol% of GM1_{extended}. For comparison, in the GM1-free control system the decay constant to describe the rate of phase separation was 0.9 μs . These data suggest phase separation in a leaflet opposite to the GM1_{short}-rich monolayer to take place somewhat faster than phase separation opposite to a GM1_{extended}-rich leaflet.

Similar conclusions can be drawn from Figure 8. It shows how 6 mol% of GM1_{extended} slows down membrane registration significantly. More importantly, Figure 8 highlights that in the system with 6 mol% of this long-chain GM1 (GM1_{extended}), the interdigitating GM1 chains have a clear effect on the opposing leaflet, and disturbing its phase behavior (see Fig. 7) leads to the weakest domain registration between the two leaflets among the systems we studied.

Discussion

Glycosphingolipids constitute a diverse class of molecules both in terms of their glycan structure and fatty acids. Most frequently, fatty acids in GSLs are C16, C18, C20, C22, C24, and C24:1 [80], though shorter [81] C12 and C14 and longer [82, 83] C26-C36 fatty acids occur. Experimental evidence suggests that the chain length can affect ligand properties of the glycan headgroup. For instance, it was shown that long-chain GSLs enhance the binding of verotoxin [81]. Galectin-4, involved in apical transport, has been observed to have a strong preference for long-chain sulfatide (C24) [52]. In our study, we have selected GM1, known for its counterreceptor status for bacterial and tissue lectins. Earlier monolayer studies suggested that GSLs' acyl chain structure would affect the glycan's presentation, such that increasing exposure of GSL carbohydrates to the solvent would correlate positively with increasing acyl chain length [51]. However, our simulation results reported in this paper show hardly any impact of acyl chain length (ranging from 16 to 30 carbon atoms) on solvent accessibility or the conformation of the carbohydrate headgroup of GM1.

We found that a long acyl chain exhibits a very strong tendency to interdigitate, *i.e.*, to protrude deep into the opposing bilayer. A deeper penetration of these chains into the opposing bilayer leaflet was observed to correlate with a substantial increase in the ordering at the chain end. The stronger the interdigitation, the more ordered the chain ends were observed to become. The observed interdigitation that takes place in a chain-length-dependent manner establishes a coupling between the two bilayer leaflets, as lipids in one leaflet can affect the structural properties of the other. Such behavior can be general for sphingolipids that are typically characterized by an inherent length asymmetry between their two chains [84].

The presented atomistic simulation data suggest that long-chain GM1 in the extracellular leaflet could modulate the phase behavior in the opposing cytosolic leaflet through GM1-induced membrane coupling. To explore this possibility, we used coarse-grained Martini simulations that allow considerations of sufficiently long times and large system sizes. In the simulations, GM1 resides exclusively in the extracellular leaflet, fully in line with experimental data. Our results showed spontaneous phase separation in GM1-containing membranes, where GM1 preferentially segregated into L_o domains. Given that the GM1 concentration in the simulations was low, the observed separation is in line with earlier Martini simulation studies [85]. Furthermore, based on experiments, phase separation takes place in mixtures of PSM and cholesterol together with dioleoylphosphatidylcholine [86]. In the present simulations for GM1 with a shorter acyl chain (~ 18 carbons), the two bilayer leaflets exhibited perfect registry, *i.e.*, they were phase-symmetric. In contrast, GM1 with an extended acyl chain (~ 30 carbons) was observed to perturb the phase behavior of the GM1-free cytosolic leaflet. A long GM1 acyl chain that interdigitated into the opposite leaflet induced mixing between saturated and unsaturated lipids, thereby preventing or at least slowing down the ideal phase separation in the cytosolic leaflet. This result is consistent with an earlier experimental study, which showed the existence of an intermediate phase in the region opposite to an L_o domain [23].

In aggregate, the present study shows a connection between GM1 acyl chain length and bilayer phase behavior. Interleaflet coupling is strongly promoted by long-chain GM1-induced interdigitation, which modulates the phase behavior in the opposing membrane leaflet. Such a physical interaction between the leaflets offers a plausible mechanism for mediating or fostering signaling across asymmetric membranes.

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Supplementary Information

Additional results and discussion.

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Table 1. Compositions of the atomistic and coarse-grained lipid bilayer models.

Bilayer	Lipid composition	GM1 type (number of carbon atoms or beads in the acyl chain)	Simulation length
Symmetric bilayers (all-atom model)	GM1:Chol:POPC = 4.86:25:70.14	GM1 ₁₆ (16 carbon atoms) or GM1 ₁₈ (18 carbon atoms) or ... or GM1 ₃₀ (30 carbon atoms)	400 ns each (16 bilayers) = 6.4 μ s
	GM1:POPC = 6.48:93.52		
Asymmetric bilayers (coarse- grained model)	GM1:PSM:DLiPC:Chol = 1.50:28.5:40:30 [GM1-rich leaflet] GM1:PSM:DLiPC:Chol = 0:30:40:30 [GM1-free leaflet]	GM1 _{short} (5 beads, corresponds to ~18 carbon atoms) or GM1 _{extended} (8 beads, corresponds to ~30 carbon atoms)	For 1.5 mol%, 10 μ s each (3 bilayers) x 10 repeats each = 300 μ s; for 6 mol% GM1 _{extended} , 20 μ s x 20 repeats = 200 μ s.
	GM1:PSM:DLiPC:Chol = 6:27:38.5:28.5 [GM1-rich leaflet] GM1:PSM:DLiPC:Chol = 0:30:40:30 [GM1-free leaflet]		
Symmetric bilayer (coarse- grained model)	GM1:PSM:DLiPC:Chol = 0:30:40:30	GM1-free model (control system)	10 μ s x 8 repeats = 80 μ s

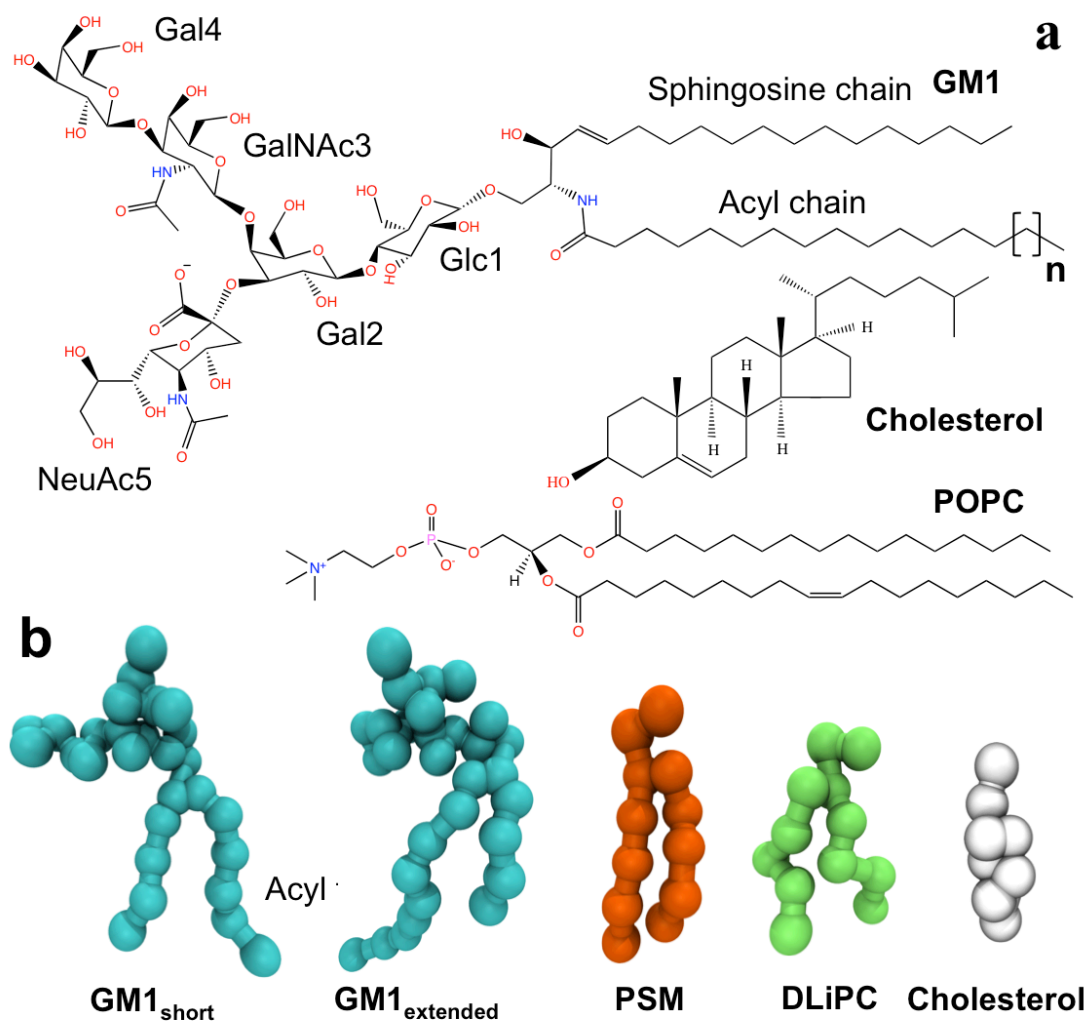


Figure 1. Chemical structure different lipid species used for (a) all-atom and (b) coarse-grained simulations. In atomistic simulations, the acyl chain length of GM1 was varied from 16 to 30 carbon atoms. In coarse-grained simulations, GM1 had an acyl chain with either five (GM1_{short}) or eight beads (GM1_{extended}).

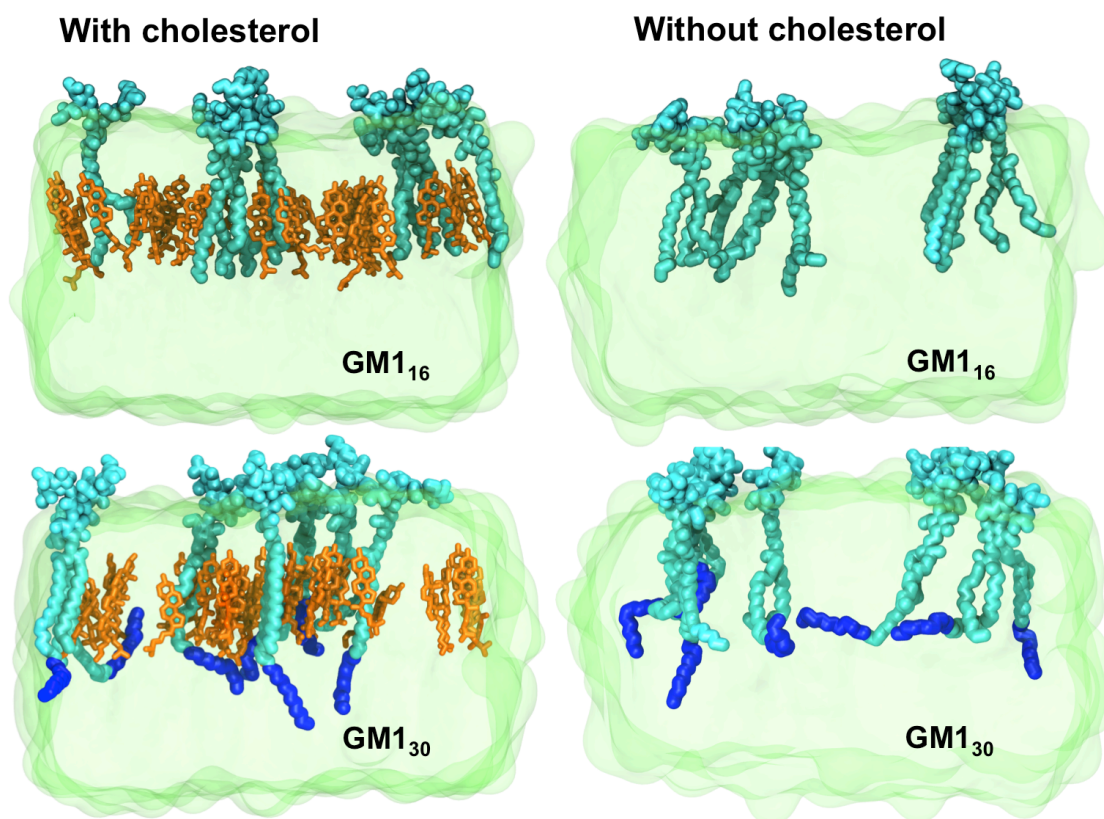


Figure 2. Snapshots of symmetric bilayers with the shortest (GM1₁₆, upper panel) and the longest (GM1₃₀, lower panel) ganglioside in the atomistic simulations. For clarity, bilayers are shown as a green transparent surface, where dark green depicts the edges of the bilayer region. GM1 (cyan) and cholesterol (orange) molecules are shown only in the upper leaflet, and POPC is not shown. For GM1₃₀, carbon atoms beyond the 18th carbon are shown in blue.

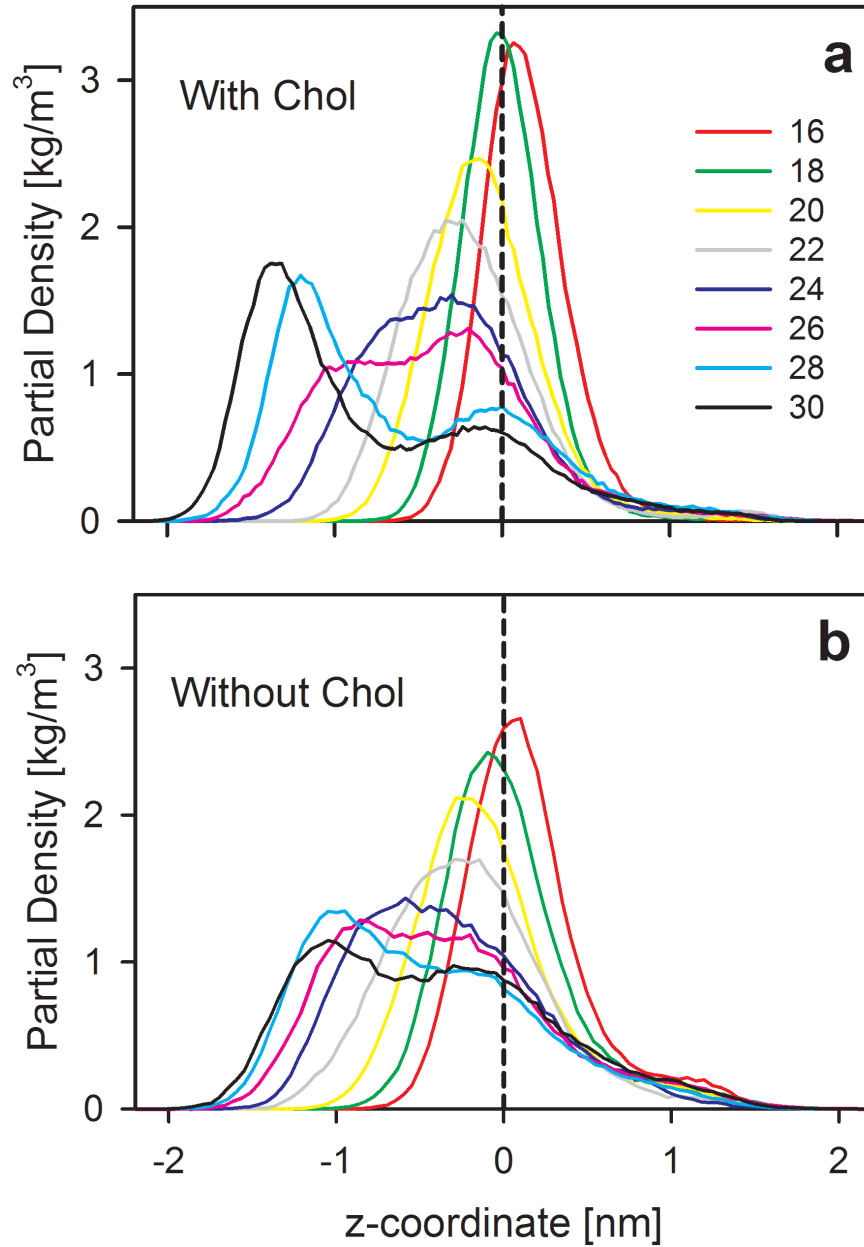


Figure 3. Partial density profiles (atomistic simulations) showing the distribution of the terminal carbon atom of the GM1 acyl chain in membranes (a) with cholesterol (Chol) and (b) without it, with varying length of the GM1 acyl chain. GM1 molecules considered here locate in the upper leaflet with $z > 0$ and interdigitate to the lower leaflet with $z < 0$.

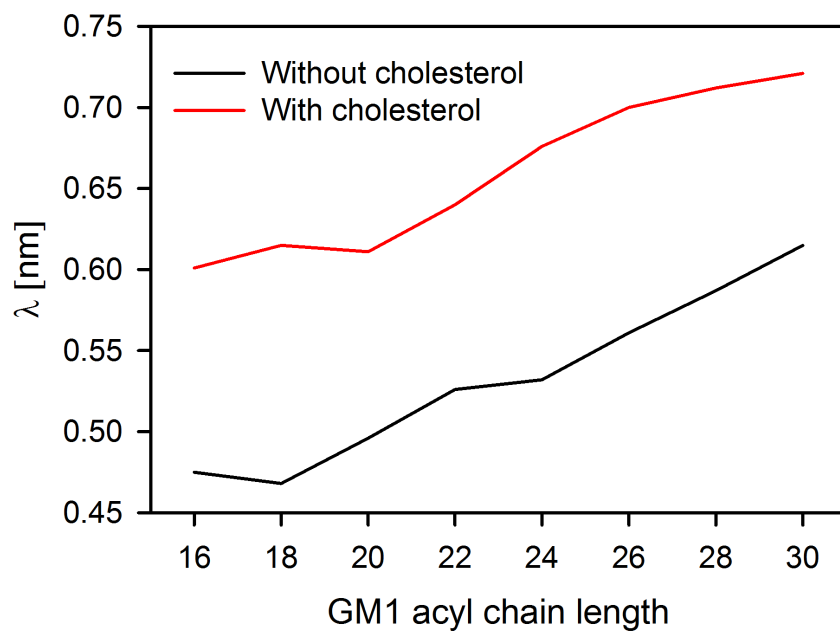


Figure 4. Values of λ (in units of nm) for systems (red) with and (black) without cholesterol as a function of GM1 acyl chain length (atomistic simulations).

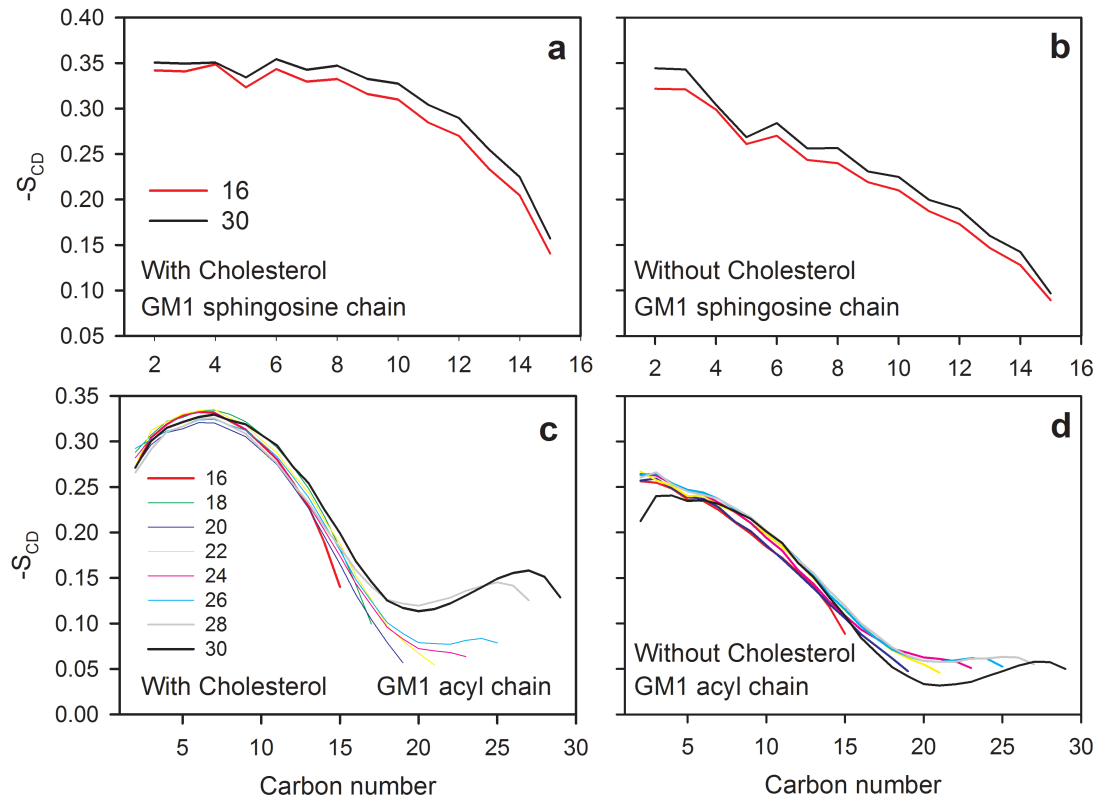


Figure 5. S_{CD} order parameter profiles of GM1 chains shown in terms of the carbon number in the chain: (a, b) sphingosine and (c, d) acyl chain, in membranes (a, c) with, and (b, d) without cholesterol.

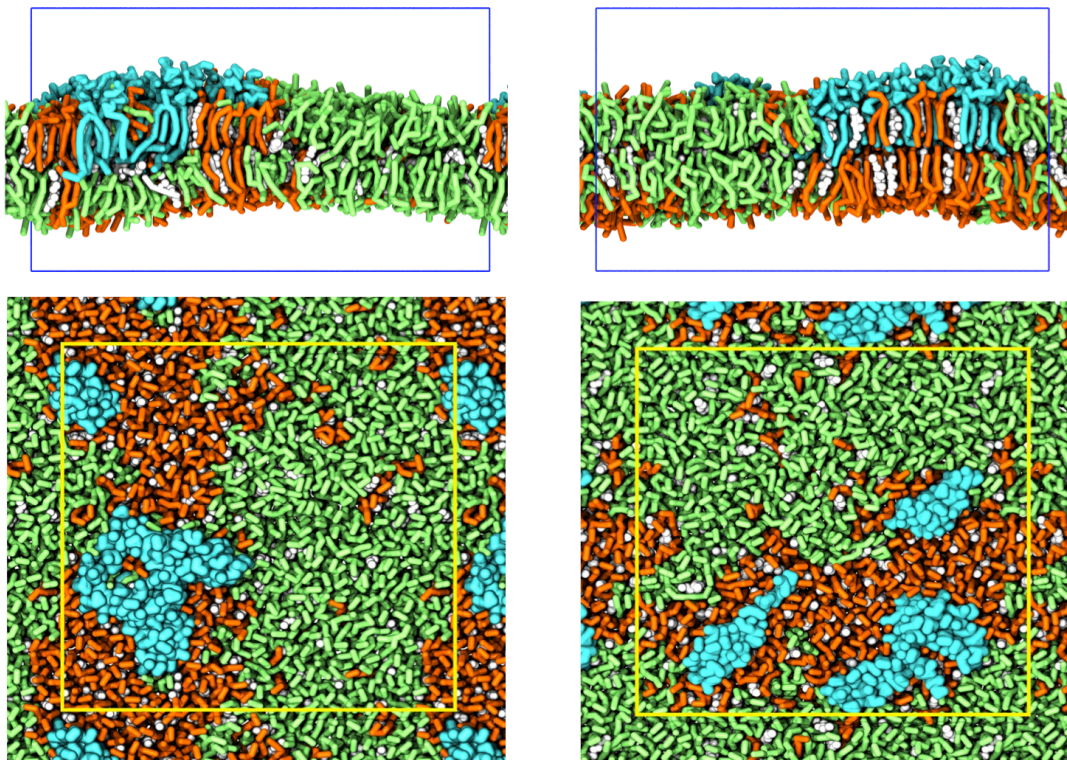


Figure 6. Final simulation structures of coarse-grained systems containing 6 mol% of (left) GM1_{extended} or (right) GM1_{short}, viewed from (top row) the side and from (bottom row) above. PSM is pictured in orange, DLiPC in green, cholesterol in white, and GM1 in cyan. Water and ions have been omitted from the images for clarity.

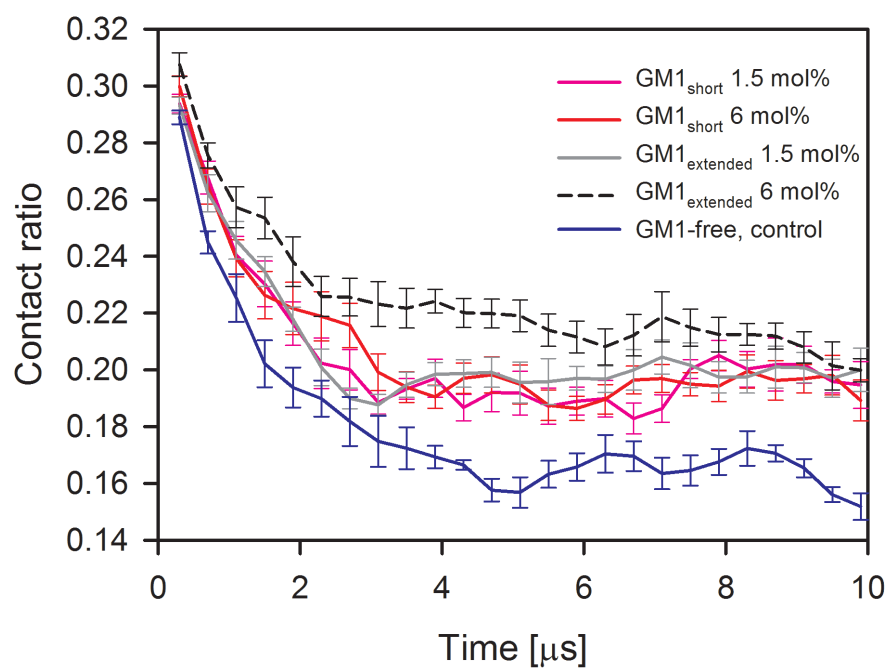


Figure 7. Contact ratio between PSM and DLiPC lipids during CG simulations that start from a random initial configuration and evolve towards phase separation.

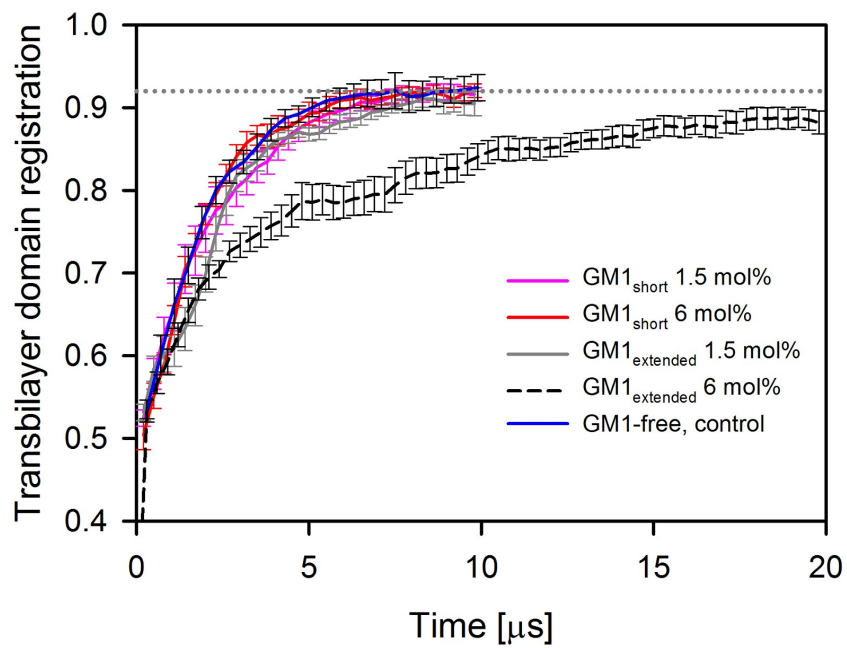


Figure 8. Transbilayer domain registration in the different systems (CG simulations). The gray dotted line highlights the value where the data of the simulated systems (except for GM1_{extended} (6 mol%)) converge.

Table of Contents Graphic

